Applicant:
 Rosenbluth et al.
 PATENT

 Serial No.:
 10/726,135
 Atty Docket: 388700-029G

Art Unit: 3731

## **AMENDMENTS TO THE CLAIMS**

Please amend claims 79, 90 and 117 as set forth below.

Please cancel claims 107-110 as set forth below

Please add new claims 118-128 as set forth below.

## Listing of Claims

## 1-78. (Canceled)

- 79. (Currently Amended) A method for preventing leakage into a perigraft space between an endovascular graft that has been implanted in the lumen of a blood vessel of a human or veterinary patient and an adjacent portion of the blood vessel wall, said method comprising the steps of:
  - (A) providing a device comprising a solid member having expansile polymeric material disposed thereon, said expansile polymeric material being i) initially in a non-expanded state wherein a quantity of the polymeric material occupies a first volume and b) subsequently expandable to an expanded state wherein said quantity of the polymeric material occupies a second volume larger than the first volume and absorbs blood;
  - (B) inserting a cannula into <u>said lumen of the blood vessel</u>; a <u>perigraft space</u> between the endovascular graft and the blood vessel wall;
  - (C) disposing said endovascular graft over a distal end of said cannula and over said adjacent portion of said blood vessel wall;
  - (D) introducing the <u>said</u> device through the <u>said</u> cannula and into <u>a</u> the perigraft space <u>between said endovascular graft and said blood vessel wall</u> while <u>said</u> the expansile polymeric material is substantially in its non-expanded state;

 Applicant:
 Rosenbluth et al.
 PATENT

 Serial No.:
 10/726,135
 Atty Docket: 388700-029G

Art Unit: 3731

monomers

(E) allowing the <u>said polymeric</u> material to expand to its expanded stated within the <u>said perigraft</u> space such that the <u>said device</u> substantially fills the

said perigraft space.

80. (Previously Presented) A method according to claim 79 wherein i) the adjacent portion of the blood vessel wall is aneurysmic; ii) the endovascular graft is implanted within the blood vessel such that it extends through the aneurysmic portion of the blood vessel and defines a perigraft space between the graft and the aneurysmic wall of the blood vessel; and, iii) the device is introduced into the perigraft space where the

expansile polymeric material expands to substantially fill the perigraft space.

81. (Currently Amended) A method according to claim 80 wherein the total volume of non-expanded expansile polymeric material introduced in Step & D is predetermined to substantially fill the perioraft space after it has been allowed to expand in Step E D.

82. (Previously Presented) A method according to claim 79 wherein the expansile polymeric material is radiopaque.

83. (Previously Presented) A method according to claim 82 wherein the expansile polymeric material is rendered radiopaque by the incorporation of radiopaque

84. (Previously Presented) A method according to claim 79 wherein the polymeric material expands to its expanded state when the pH of its environment is a physiological pH of about 7.4.

85. (Previously Presented) A method according to claim 79 wherein the polymeric material is in the form of pellets when introduced through the cannula.

Applicant: Rosenbluth et al. PATENT
Serial No.: 10/726.135 Atty Docket: 388700-029G

Art Unit: 3731

86. (Previously Presented) A method according to claim 79 wherein the solid

member is an elongate member.

87. (Previously Presented) A method according to claim 86 wherein the solid

member is filamentous.

88. (Previously Presented) A method according to claim 86 wherein a plurality of

pieces of the polymeric material are disposed at spaced-apart locations on said

elongate solid member.

89. (Previously Presented) A method according to claim 88 wherein the device

further comprises coil spacers disposed on said solid member between pieces of the

expansile polymeric material.

90. (Currently Amended) A method according to claim 79 wherein the solid member

is formed of platinum platmium.

91. (Previously Presented) A method according to claim 79 wherein the solid

member is formed of platinum and tungsten.

92. (Previously Presented) A method according to claim 79 wherein the solid

member is formed of wire.

93. (Previously Presented) A method according to claim 79 wherein the solid

member is formed of polymeric material.

94. (Previously Presented) A method according to claim 93 wherein the solid

member is formed of a polymer filament.

95. (Previously Presented) A method according to claim 94 wherein the solid

member is formed of a polyvinyl alcohol filament.

Applicant: Rosenbluth et al. PATENT
Serial No.: 10/726.135 Atty Docket: 388700-029G

Art Unit: 3731

96. (Previously Presented) A method according to claim 79 wherein the solid

member is biased to a coiled configuration.

97. (Previously Presented) A method according to claim 79 wherein the cannula is

advanced through the lumen of a catheter.

98. (Previously Presented) A method according to claim 97 wherein the catheter is a

microcatheter.

99. (Previously Presented) A method according to claim 98 wherein the

microcatheter has a lumen of 0.005-0.050 inch diameter.

100. (Previously Presented) A method according to claim 79 wherein the device is

initially attached to a delivery member by way of a detachable connection, said delivery member being useable to advance the device into the perigraft space, said detachable

connection being thereafter detachable such that the delivery member may be retracted

into the cannula while the device remains in the perigraft space.

101. (Previously Presented) A method according to claim 79 wherein the polymeric

material expands more rapidly as the pH of its environment increases.

102. (Previously Presented) A method according to claim 79 wherein the polymeric

material is a hydrogel.

103. (Previously Presented) A method according to claim 79 wherein the polymeric

material is porous when in its expanded state.

104. (Previously Presented) A method according to claim 103 wherein the porous

polymeric material, when substantially fully expanded, has pores of about 50-1000

microns in diameter

Applicant: Rosenbluth et al. PATENT
Serial No.: 10/726.135 Atty Docket: 388700-029G

Art Unit: 3731

105. (Previously Presented) A method according to claim 103 wherein the porosity of

the polymeric material, when substantially fully expanded, is at least about 50%.

106. (Previously Presented) A method according to claim 103 wherein the porosity of

the polymeric material, when substantially fully expanded, is between about 50% and

about 95%.

107-110. (Canceled)

111. (Previously Presented) A method according to claim 79 wherein the cannula is

substantially rigid.

112. (Previously Presented) A method according to claim 79 wherein the cannula is

substantially flexible.

113. (Previously Presented) A method according to claim 79 wherein the cannula

comprises a metal tube.

114. (Previously Presented) A method according to claim 79 wherein the cannula

comprises a plastic tube.

115. (Previously Presented) A method according to claim 79 wherein the method is

performed after an endoleak has been detected as a means of treating the endoleak.

116. (Previously Presented) A method according to claim 79 wherein the method is

performed before an endoleak has been detected as a means for preventing an

endoleak from occurring.

117. (Currently Amended) A method according to claim 79 wherein Step B

comprises: advancing a catheter to a first position within the patient's vasculature; and,

Page 6 of 13 13468\_1.DOC

 Applicant:
 Rosenbluth et al.
 PATENT

 Serial No.:
 10/726.135
 Atty Docket: 388700-029G

Art Unit: 3731

advancing the cannula through the catheter to a second position—wherein the distal end of the cannula is within the perigraft space.

118. (New) A method of treating a vessel within a body comprising:

positioning a distal end of a delivery device in proximity of a target location within said vessel:

expanding a graft at said target location within said vessel such that a perigraft space is formed, said graft expanding over said distal end of said delivery device; and

introducing an expansile material into said perigraft space through said delivery device;

expanding said expansile material within said perigraft space.

119. (New) The method of claim 118, wherein said positioning a distal end of a delivery device in proximity of a target location within said vessel further comprises positioning a cannula adjacent an aneurysmic wall.

120. (New) The method of claim 119, wherein said expansile material is disposed on a solid member.

121. (New) The method of claim 119, wherein said expansile material is disposed on a coil.

122. (New) The method of claim 119, wherein said expansile material is a hydrogel.

123. (New) The method of claim 119, wherein said expansile material expands at a pH of about 7.4.

124. (New) A method of treating an aneurysm of a vessel comprising: advancing a distal end of a cannula to an aneurysm; 
 Applicant:
 Rosenbluth et al.
 PATENT

 Serial No.:
 10/726,135
 Atty Docket: 388700-029G

Art Unit: 3731

positioning an endovascular graft near said aneurysm;

anchoring said endovascular graft to a wall of the vessel over said aneurysm;

capturing said cannula between said graft and said vessel wall:

delivering an expansile material into said aneurysm through said cannula.

125. (New) The method of claim 124, wherein said advancing a distal end of a cannula to an aneurysm further comprises advancing a catheter to a position in

proximity to the aneurysm.

126. (New) The method of claim 124, wherein said delivering an expansile material into said aneurysm through said cannula further comprises delivering a hydrogel into

the perigraft space.

127. (New) The method of claim 124, wherein said delivering an expansile material into said aneurysm through said cannula further comprises delivering the expansile

material disposed on a coil.

128. (New) The method of claim 124, wherein said delivering an expansile material into said aneurysm through said cannula further comprises delivering a pH sensitive

hydrogel.